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# Molecular analysis and associated pathology of beak and feather disease virus isolated in Italy from young Congo African grey parrots (*Psittacus erithacus*) with an "atypical peracute form" of the disease

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## Abstract

This study is the first report on the genetic and pathogenic characterization of beak and feather disease virus (BFDV) occurring in Italy. Twenty BFDV strains isolated in Italy from juvenile Congo African grey parrots (*Psittacus erithacus*) were investigated. Seventeen strains showed an "atypical peracute form" (aPF) of the disease, and three a chronic form (CF). Usually, the peracute form affected neonatal or fledgling chicks, presenting with haemorrhage and/or multiorgan necrosis. The affected birds had been weaned, were independent as far as food and protection were concerned and apparently were without lesions. The gene coding for the putative coat protein was amplified in all isolates while the BFDV genome was sequenced completely in 10 samples, eight of them belonging to aPF affected birds and two from CF of the disease. All full genomes clustered into the J strain of BFDV, where two new subtypes were identified. Recombination analyses showed evidence of genetic exchanges in two BFDV genomes. In addition, a correlation between viral isolate and origin of the breeding material was shown, while an association between the genetic features of the virus and the clinical form was not observed. Histologically, apoptosis was detected frequently in aPF samples and sporadically in CF samples. Interestingly, BFDV antigens were detected in the nuclei and cytoplasm of such apoptotic cells. The data presented here support the hypothesis that, in the absence of a defined BFDV genetic variant accountable for a specific clinical form of psittacine beak and feather disease, differences in the apoptotic rate among aPF and CF are strictly host related.

**Keywords:** beak and feather disease virus, *Psittacus erithacus*, phylogenetic analysis, Italy

## Introduction

Psittacine beak and feather disease (PBFD) is a cosmopolitan disease affecting numerous species of domestic and wild birds of the order Psittaciformes in the world. The disease is caused by a small virus roughly 2 kb in size characterized by a non-enveloped, icosahedral, circular, ambisense, non-segmented, single-stranded DNA. Beak and feather disease virus (BFDV) is a member of the genus *Circovirus*, family *Circoviridae*. These viruses are composed of up to seven open reading frames (ORF1 to ORF7), two of which, ORF1 and the ORF2, are well described, while the others are not always present and are less characterized (Ritchie *et al.*, 1989; Bassami *et al.*, 1998; Heath *et al.*, 2004; Julian *et al.*, 2013). ORF1 (*rep* gene) is present in the virion sense strand, encodes the viral replication-associated protein Rep (Bassami *et al.*, 1998; Faurez *et al.*, 2009) and shows highly conserved amino acid motifs (Mankertz *et al.*, 1998; Heath *et al.*, 2004). ORF2 (*cp* gene) translated in the complementary strand encodes the major structural capsid protein (CP), responsible for the encapsidation of the virus and its entry into the cells as well as for the viral crossing of the nuclear envelope (Ritchie *et al.*, 1989; Niagro *et al.*, 1998; Heath *et al.*, 2006). These two ORFs display different rates of evolution, with the Rep being relatively more conserved than the CP (Varsani *et al.*, 2011; Julian *et al.*, 2013). The intergenic region between the two major ORFs contains a replication hairpin loop structure (TAGTATTAC) that is characteristic and highly conserved in all BFDV circo viruses. These viruses have been described as ecologically stable and can be transmitted

by both horizontal and vertical routes (Ritchie, 1995; Niagro *et al.*, 1998; Rahaus *et al.*, 2008). Usually, the clinical forms vary from species to species and depend on the age of the bird when infected (Gerlach, 2004). PBFD is a potentially fatal disease and can be present as a peracute, acute, or chronic form (CF) of infection; the latter is the most frequent form, known as the "classical form". Frequently, in birds up to 3 years of age, the disease begins with depression and lethargy followed by a chronic, progressive, bilateral and symmetrical feather dystrophy occasionally accompanied by beak deformity. Until now only peracute and acute forms have been described in nestling and fledgling parrots (Schoemaker *et al.*, 2000; Doneley, 2003; Rahaus & Wolff, 2003; Raue *et al.*, 2004). Here, death may occur suddenly, without evidence of feather dystrophy (peracute form) or with mild evidence of feather dystrophy (acute form). In addition, BFDV-infected birds, especially those with chronic infection, typically display lymphoid depletion and immunosuppression that favour the appearance of opportunistic secondary infections (Ritchie *et al.*, 1989; Todd, 2000), often leading to death. Currently, there is no cure or effective treatment for the disease caused by BFDV. Moreover, the production of a protective vaccine is hampered by the lack of success in culturing BFDV *in vitro*.

The international trade in exotic parrots, both through legal trade and illegal trafficking, has facilitated the spread of BFDV, such that BFDV can now be considered a global disease. It is likely that in Italy, as in Poland (Julian *et al.*, 2013), there have been multiple introductions of BFDV over a long period of time. Consequently, captive infected birds kept in close proximity have favoured the recombination of different BFDV strains leading to the evolution of new viral subtypes. An epidemiological study carried out in Italy (Bert *et al.*, 2005) showed that there was a high occurrence of infection in parrots imported from the "Old World" (South East Asia, Africa). In these birds the infection was mainly restricted to Cacatua and Congo African grey parrots (CAGPs), indicating a high susceptibility to BFDV in these two species. Over the years, several classifications have been proposed by different authors (Ritchie *et al.*, 2003; de Kloet & de Kloet, 2004; Heath *et al.*, 2004; Varsani *et al.*, 2011). Following the recent classification system proposed by Varsani *et al.* (2011), the predominant strain circulating in Europe is BFDV-J, a strain isolated mainly from *Psittacus erithacus*. Other strains circulating in these parrots in Europe are BFDV-I (Poland and Portugal), BFDV-J (Germany, Poland, Portugal, the UK) and BFDV-T (Poland), as reported by Varsani *et al.* (2011) and Julian *et al.* (2013).

The purpose of this study was to characterize the BFDV strains circulating amongst juvenile CAGPs that died of an "atypical peracute form" (aPF) of the disease, involving weaned parrots 4 to 6 months of age, and to document the associated histopathology. Finally, this study looked at the phylogenetic relationships of these strains in comparison with previously reported sequences.

## Materials and Methods

**Study design.** Between April 2008 and November 2011, all 4-month-old to 6-month-old hand-reared, weaned and fully fledged CAGPs from well-managed breeding centres in Northern and Central Italy that died suddenly were submitted to our laboratory for investigation. Before death, some of the birds had displayed mild lethargy and anorexia. The occurrence of BFDV was confirmed by polymerase chain reaction (PCR), using the protocol described by Ypelaar *et al.* (1999). The presence of other pathogens, such as *Chlamydia psittaci*, Polyoma virus and herpesvirus (Pacheco's disease), which could present a similar clinical picture, were excluded by molecular analysis (PCR amplification). Prior to this investigation, parents of infected birds and other randomly selected parrots in each breeding centre had been tested. All centres were negative for BFDV with the exception of one, where we documented a 2% incidence of BFDV in asymptomatic birds.

**Sample collection.** This study was carried out on samples of various organs from 17 young CAGPs (4 to 6 months of age) that died suddenly with an aPF of BFDV. Samples from three adult birds (3, 8, and 11 years old respectively) that had died with typical signs of the CF of BFDV (such as abnormal feathers and beak lesions) were also included. The samples from birds with aPF originated from nine breeding centres: seven from one centre, three from another centre and one sample each from the remaining seven centres. The samples from the adult birds were provided by three private owners (Table 1).

**Histology, immunohistochemistry and TUNEL evaluation.** For 10 of the 20 CAGPs analysed in the study (eight showing an aPF and two a CF of the disease), a full necropsy was performed and target organs were sampled for histological, immunohistochemical and terminal

deoxynucleotidyl transferase dUTP nick end-labelling (TUNEL) examination. The organs analysed were the liver, spleen, thymus, heart, lung, kidney, bursa of Fabricius and femoral bone marrow (BM). These samples were fixed in 10% neutral buffered formalin, paraffin embedded and serially sectioned at 3  $\mu$ m.

Archived formalin-fixed, paraffin wax-embedded tissues from three CAGPs with no history of clinical diseases and negative for BFDV on PCR were retrieved from the University of Camerino Veterinary Pathology Unit archives. These samples had been obtained at post-mortem from parrots that were presented for euthanasia following severe trauma and their ages ranged from 8 to 11 years. These samples were used as histological controls.

#### *Immunohistochemical tests*

For this purpose, rehydrated sections were treated for endogenous peroxidase neutralization with 3% hydrogen peroxide for 5 min followed by rinsing for 5 min in distilled water. Antigen retrieval was achieved by incubating the slides in two antigen retrieval solutions—citrate buffer, pH 6.0 (for BFDV and interleukin [IL]-1 $\beta$ ); and 0.01 M Tris-ethylenediamine tetraacetic acid buffer, pH 9.0 (for caspase-1)—in a steamer according to the instructions of the manufacturer (Black & Decker, Towson, MD, USA) for 20 min. Non-specific immunoglobulin binding was blocked by incubating the slides for 10 min with a protein-blocking agent (Dako, Carpinteria, CA, USA). The slides were then incubated overnight in a moist chamber with the following primary antibodies: anti-BFDV hyper-immune serum (from an adult CAGP with CF of the disease, diagnosed by PCR, and highly reactive with intracellular inclusions of BFDV within affected tissues), rabbit anti-caspase-1 polyclonal antibody (Cell Signaling Technology, San Diego, CA, USA), and rabbit anti-chicken IL-1 $\beta$  (ACRIS Antibodies, San Diego, CA, USA). In brief, parrot serum, diluted 1:50 in phosphate-buffered saline+bovine serum albumin, anti-caspase-1 and anti IL-1 $\beta$  antibodies, both diluted 1:100 in the same buffer, were added and incubated overnight. Then, following three washing steps, biotinylated rabbit anti-chicken (Nordic Immunology, Tilburg, the Netherlands) or biotinylated goat anti-rabbit (Dako) antibodies were used.

Positive cells were detected with a streptavidin—immunoperoxidase staining procedure (Dako) using 3,3'-diaminobenzidine as the substrate (Vector, Burlingame, CA, USA). Tissue sections were counterstained with Mayer's haematoxylin. Immunohistochemical staining controls were slides in which primary antibodies were omitted. In the BM and lymphoid organ sections, the apoptotic index was highlighted through a TUNEL colorimetric staining (DeadEnd; Promega Madison, WI, USA) according to the manufacturer's instructions. For the evaluation of the apoptotic rate, 10 random fields in the tissue section were examined under a 40 x dry objective. The number of BFDV, caspase-1 and TUNEL-positive cells was normalized to the number of cells per field and expressed as a percentage.

IL-1  $\beta$ -positive cells were quantified in different lymphoid organs and BM, using a light microscope (Carl Zeiss Oberkochen, Germany), and a 40 $\times$  objective. Ten sites were chosen and the mean value of the cellular staining with the anti-chicken IL-1 $\beta$  primary antibody was scored as follows: score 0, <25% of cells labelled; score 1, 25 to 50% of cells labelled; score 2, 50 to 75% of cells labelled; and score 3, >75% cells labelled. For all parameters, the cells on the margins of the tissue sections were not considered for evaluation.

**Molecular analysis.** The gene coding for the putative *cp* was amplified by PCR in all isolates (17 from birds with aPF and three from birds with CF). Moreover, the BFDV genome was completely sequenced in 10 samples (eight from birds with aPF and two from birds with CF). The samples were identified using a progressive number on the basis of the year of isolation, the two-letter code identifying the country and the year of isolation (Table 1). Briefly, for DNA extraction, 25 mg mixed organs stored in 100% ethanol were cut into small pieces and processed using the DNeasy Blood & Tissue Kit (Qiagen, Milan, Italy), according to the manufacturer's instructions. Purified DNA was quantified and stored at -80°C until use.

The full-length genomes of BFDV strains were amplified using several overlapping primer sets (Table 2). PCR amplifications were carried out in a total volume of 50  $\mu$ l containing 1 $\times$  storage buffer (Qiagen), 1.5 mM MgCl<sub>2</sub> (3.5 mM MgCl<sub>2</sub> only for coat protein fragment amplification), 0.2 mM of each dNTP, 0.5 pmol of each primer, 2.5 U Taq DNA polymerase (Qiagen) and 100 ng template DNA. PCR cycling conditions were as follows: 3 min denaturation at 94°C, 35 cycles with steps of 30 sec at 94°C, 30 sec at 58 or 60°C and 1 min at 72°C, followed by a final extension of 5 min at 72°C. PCR products were examined for correct size by agarose (2%) gel electrophoresis in 1 $\times$  TAE buffer, purified by ExoSAP-IT Clean-Up (USB, Affymetrix, Cleveland, OH, USA Inc.) and sequenced by MacroGen Services (Amsterdam, the Netherlands).

**Phylogenetic analysis.** Sequences obtained from positive samples were aligned with known BFDV sequences available on the GenBank database using the ClustalW program (Thompson *et al.*, 1997). More specifically, 20 *cp* sequences obtained in this study were compared with 37 *cp* sequences of strains isolated from CAGPs in other parts of the world and the full genome of the 10 strains was aligned with the complete genome available on the GenBank database of 23 strains from CAGPs (Table 1). Phylogenetic analysis on the data *rep* ( $n = 10$ ) and *cp* ( $n = 20$ ) sequences was performed. The aligned sequences were imported into the PAUP\* version 4.0b 10 program (Swofford, 2003). The model of molecular evolution was estimated using a hierarchical likelihood ratio test approach and the Akaike information criterion (Akaike, 1973) implemented in the ModelTest version 3.7 computer program (Posada & Crandall, 1998; Posada, 2001). Bayesian methods implemented in the MrBayes version 3.1.1 program (Huelsenbeck & Ronquist, 2001; Ronquist & Huelsenbeck, 2003) were used to draw phylogenetic trees and assess statistical support for clades. In detail, a Markov chain Monte Carlo search for 1,000,000 generations using two runs with four chains (temperature = 0.05) was performed, and results were represented as a 50% majority rule consensus tree. The sequence identity comparisons were made by the Nei-Gojobori method (Nei & Gojobori, 1986). Recombination amongst all complete genomes from this study and complete sequences available in GenBank was analysed using the Recombination Analysis Tool (Etherington *et al.*, 2006). The potential recombination events were evaluated using the default program setting when phylogenetic evidence of recombination was observed.

**Sequence accession numbers.** The sequences were submitted to GenBank. The accession numbers of whole sequences of BFDV were KF723391 (1IT08), KF723386 (2IT09), KF723385 (3IT09), KF72389 (4IT09), KF723393 (5IT09), KF723384 (6IT09), KF723388 (7IT09), KF723387 (8IT10), KF723390 (9IT11) and KF723392 (10IT11). The accession numbers for the *cp* sequence were KF723402 (11IT09), KF723401 (12IT09), KF723395 (13IT09), KF723403 (14IT09), KF723399 (15IT10), KF723394 (16IT10), KF723400 (17IT10), KF723398 (18IT11), KF723397 (19IT11) and KF723396 (20IT11).

**Table 1.** Summary of information relating BFDV full genomes and putative coat protein genes from African grey parrots (*Psittacus erithacus*) determined in this study (A and B) and obtained by GenBank (C and D).

A. Full genomes determined in this study					B. Sequences of the coat protein gene determined in this study			
Strain	Country, Year	Clinical signs	Accession no.	subtype	Strain	Country, Year	Clinical signs	Accession no.
<b>1IT08</b>	North Italy, 2008	peracute	KF723391	J7	<b>11IT09<sup>a</sup></b>	Central Italy, 2009	peracute	KF723402
<b>2IT09<sup>a</sup></b>	Central Italy, 2009	peracute	KF723386	J1	<b>12IT09<sup>a</sup></b>	Central Italy, 2009	peracute	KF723401
<b>3IT09<sup>a</sup></b>	Central Italy, 2009	peracute	KF723385	J1	<b>13IT09<sup>a</sup></b>	Central Italy, 2009	peracute	KF723395
<b>4IT09<sup>a</sup></b>	Central Italy, 2009	peracute	KF723389	J1	<b>14IT09<sup>a</sup></b>	Central Italy, 2009	peracute	KF723403
<b>5IT09</b>	North Italy, 2009	peracute	KF723393	J1	<b>15IT10</b>	North Italy, 2010	peracute	KF723399
<b>6IT09</b>	Central Italy, 2009	peracute	KF723384	J1	<b>16IT10</b>	North Italy, 2010	peracute	KF723394
<b>7IT09</b>	North Italy, 2009	chronic	KF723388	J1	<b>17IT10</b>	Central Italy, 2010	peracute	KF723400
<b>8IT10</b>	Central Italy, 2010	chronic	KF723387	J1	<b>18IT11<sup>b</sup></b>	North Italy, 2011	peracute	KF723398
<b>9IT11</b>	North Italy, 2011	peracute	KF723390	J2	<b>19IT11<sup>b</sup></b>	North Italy, 2011	peracute	KF723397
<b>10IT11<sup>b</sup></b>	North Italy, 2011	peracute	KF723392	J6	<b>20IT11</b>	North Italy, 2011	chronic	KF723396
C. Full genomes gene obtained by GenBank					D. Sequences of the coat protein gene obtained by GenBank			
<b>PT05</b>	Portugal, 2005	classical	EU810208	I1	<b>1GE</b>	Germany	acute	AY518899
<b>PT08</b>	Portugal, 2008	aspecific	EU810207	J2	<b>2GE</b>	Germany	acute	AY518920
<b>PT08-2&amp;3</b>	Portugal, 2008	classical	GQ120621	J1	<b>7GE</b>	Germany	acute	AY518902
<b>PT09</b>	Portugal, 2009	aspecific	GQ329705	J1	<b>73GE</b>	Germany	acute	AY518914
<b>PT09-2</b>	Portugal, 2009	aspecific	GU047347	J1	<b>74GE</b>	Germany	acute	AY518915
<b>PEG07</b>	Germany, 2004	acute	AY521237	J3	<b>IND</b>	India	unknown	AY518906
<b>PEU01</b>	U.K., 2004	acute	AY521238	J1	<b>JAP</b>	Japan	unknown	AY518905
<b>PEP01</b>	Portugal, 2004	acute	AY521236	I4	<b>POR</b>	Portugal	acute	AY518901
<b>AFG3-ZA</b>	S. Africa, 2003	unknown	AY450443	I3	<b>11POR</b>	Portugal	acute	AY518912
<b>AFG4-ZA</b>	S. Africa, 2003	unknown	AY450435	C2	<b>16POR</b>	Portugal	acute	AY518910
<b>ZA-80A</b>	S. Africa, 2008	unknown	HM748920	I3	<b>PR</b>	Puerto Rico	none	AY518926
<b>ZA-55A</b>	S. Africa, 2008	unknown	HM748931	M1	<b>JB1UK</b>	United Kingdom	acute	AY518904
<b>BFDV1</b>	Thailand, 2006	unknown	GU015012	F1	<b>US34</b>	USA	acute	AY518927
<b>BFDV2</b>	Thailand, 2006	unknown	GU015013	F1	<b>US36</b>	USA	acute	AY518928
<b>PL688</b>	Poland, 2008	unknown	JX221037	J4				
<b>PL785</b>	Poland, 2009	unknown	JX221041	J5				
<b>PL336</b>	Poland, 2007	unknown	JX221018	T1				
<b>PL376</b>	Poland, 2007	unknown	JX221020	T1				
<b>PL399</b>	Poland, 2007	unknown	JX221022	T1				
<b>PL410</b>	Poland, 2007	unknown	JX221023	T1				
<b>PL573</b>	Poland, 2008	unknown	JX221032	J1				
<b>PL711</b>	Poland, 2008	unknown	JX221038	I4				
<b>PL717</b>	Poland, 2008	unknown	JX221039	I4				

<sup>a</sup> and <sup>b</sup> indicate strains obtained from birds were aviary mates, respectively.

**Table 2.** Nucleotide sequence, location (nt) of the primers, predicted sized of the amplified products, and annealing temperature of the oligonucleotide primers designed to amplify the full-length genomes of PBFDV.

Sets of primers	Primers	Sequences (5'-3')	Location <sup>a</sup> (nt)	Product size (bp)	Annealing (°C)	Reference
<b>A</b>	1F	CGAAGACTACCGRATTAAGTTA	1691-1670	579	58	This study
	2R	AACGATGGCATAGTAGAATTCG	250-229			
<b>B</b>	3F	ATGCCGTCCAAGGAGGGCT	131-149	767	58	This study
	4R	GTCACAGTCCTCCTTGATACCA	898-878			
<b>C</b>	5F	CCTTATTGCGAGATGCTCCG	758-777	638	58	This study
	6R	TGGATCCCACTACAAGGAGGA	1376-1396			
<b>D</b>	7F	ACCTCTAACTGCGCATGC	1975-1958	801	60	This study
	8R	CACGGTAGCGCCGAAGG	1174-1190			

<sup>a</sup> The position of the primers is referred to the sequence PEU01 (Acc. AY521238).

## Results

**Necropsy.** In the eight juvenile parrots that died with the aPF of BFDV, gross lesions were absent. The parrots displayed good muscle and plumage development; however, it was of interest that all of these birds were severely anaemic (based on laboratory tests performed *in vivo* by various practitioners). In comparison, the two adults with the CF of the illness showed mainly feather abnormalities such as shortness, clubbing and anomalous curling. There was also moderate atrophy of the pectoral muscles, and the livers were enlarged and mottled with scattered foci of necrosis, as were the spleens. Botryoid inclusion bodies were seen in macrophages of some of the feather follicles and in some spleen and intestinal macrophages. No abnormalities were detected in the BM of the CF group.

**Histology.** The histological picture from the aPF affected birds was similar, consisting of lymphoid depletion in the spleen, thymus and bursal follicles. There were also foci of hepatic necrosis and severe BM atrophy. The lymphoid tissue was markedly atrophic with large numbers of cells undergoing apoptosis together with scattered areas of necrosis. In some cases, the only histologic lesion in birds affected with the aPF of the disease was severe necrosis of bursal, thymic, and BM elements. In the lymphoid organs, and especially in the BM, apart from the apoptotic lymphocytes, many infiltrating macrophages were also observed (Figure 1a,b). In these birds, since viral inclusions were not visible in any of the damaged tissues (Figure 1c,d), even if the signs were suggestive of a viral infection, BFDV was not the prime suspect. The histological picture in the two parrots affected by the CF of the disease was characterized by similar lesions in the beak and feathers and included necrosis and hyperplasia of the epithelial cells in the basal and intermediate epithelial layers. The feather pulp was infiltrated mainly by heterophils, plasma cells and macrophages with lesser numbers of lymphocytes. In this group of birds, basophilic nuclear and cytoplasmic inclusions were detected in epithelial cells, whereas inclusions were frequently found in macrophages in the epidermis, feather pulp cavity and lymphoid tissue cytoplasmic (data not shown).

**Immunohistochemical and TUNEL evaluations.** Higher numbers of apoptotic cells were detected in the spleen, thymus, bursa and BM of aPF affected compared with CF affected CAGPs, by concurrent positive TUNEL assay and immunostaining for activated caspase-1 (Figure 1e,f,g,h). A large number of the caspase-1-positive cells exhibited a concomitant staining with BFDV antiserum (Figure 1c,d,e,f). Apoptotic cells were frequently detected in the enlarged germinal centres of spleens and bursa in the aPF and only sporadically in the same organs of the CF. No BFDV-positive cells and only low levels of caspase-1-positive cells were detected in the tissues from the euthanized parrots, retained as control (data not shown). The apoptotic index of the different organs in the three groups of birds (aPF, CF and euthanized) is reported in Table 3, in association with the mean percentage value of BFDV-positive cells. Apoptotic cells, identified by activated caspase-1 and TUNEL corresponded to cells with microscopic changes typical of apoptosis in haematoxylin and eosin stained sections. The cytoplasm and less frequently the nuclei of many apoptotic cells in the bursal and spleen

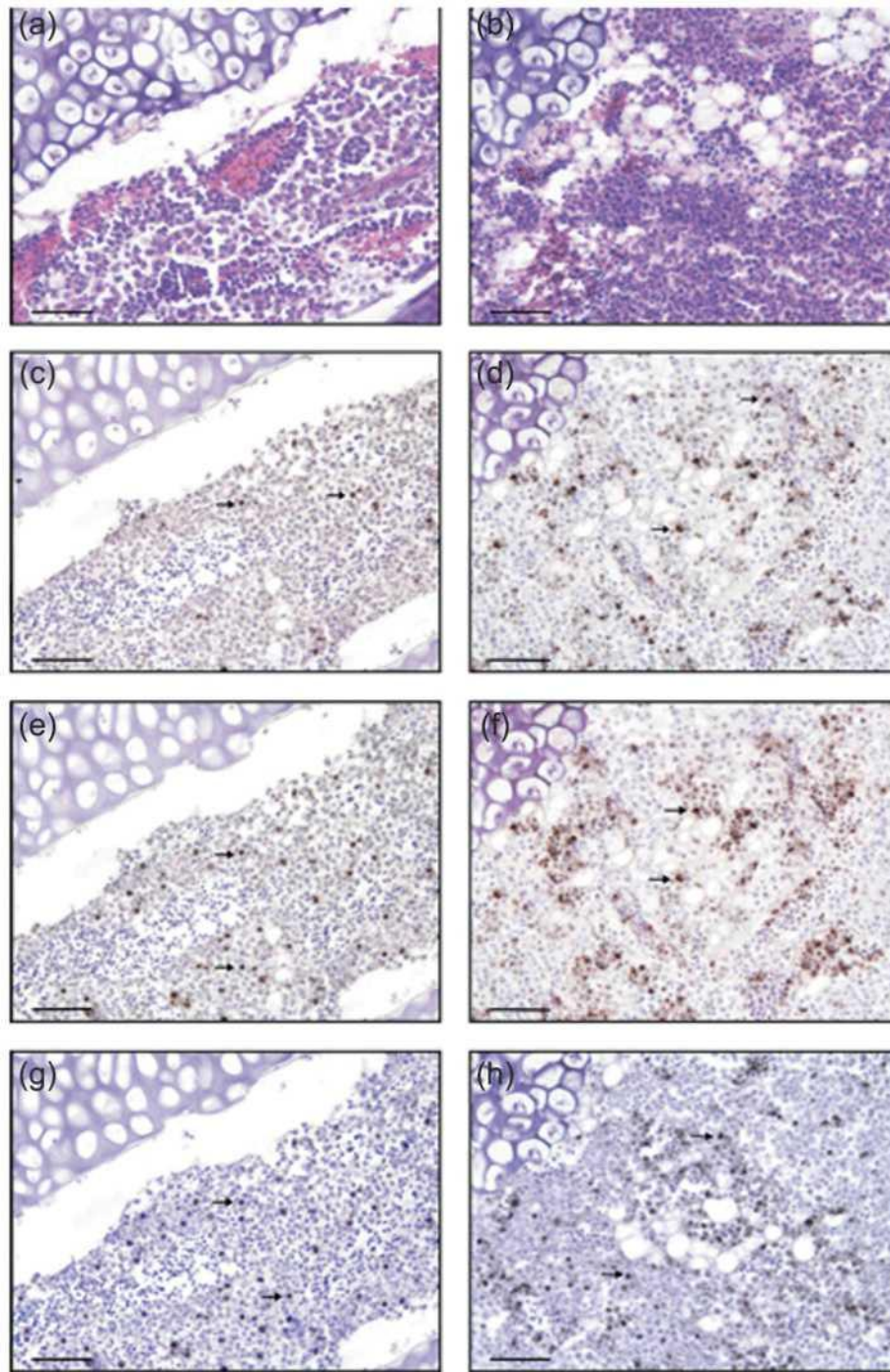


germinal centres were positive for BFDV antigens while BFDV was occasionally identified in normal and apoptotic BM cells of CF affected CAGPs (Figure 1c).

**Molecular analysis.** The complete genome sequences of 10 BFDV strains as well as nucleotide sequences of the other 10 putative *cp* genes were determined and analysed. The length of the full genomes ranged from 1997 to 2001 nucleotides with pairwise identities of between 93.6% and 99.9%. The nucleotide identity among the 20 *cp* sequences and the 10 *rep* sequences varied respectively from 88.2% to 100% and from 95.5% to 100%. The pairwise amino acid identity for *cp* and *rep* protein was >88.5% for *cp* and >98.8% for *rep*.

The relevant genomic features of the full-length Italian strains are reported in Table 4. Briefly, all genomes contained the distinguishing nonanucleotide circovirus origin of replication (TAGTATTAC) located within a stem loop structure, the octamer sequences that most probably act as a binding site for BFDV Rep protein, the TATA boxes and the polyadenylation signals. The putative ORFs varied from four to six, instead of the seven ORFs described by Bassami *et al.* (1998). All strains contained the two major ORFs: ORF1, which codes for the *rep* gene, was 870 nucleotides in length, had a canonical ATG sequence as start codon and TAA or TGA as stop codon; and ORF2, which codes for *cp* gene, was 750 nucleotides in length. The differences in the length of this gene in comparison with the data already published in the literature depend on the start codon that is chosen, since a canonical starting site is not evident (Bassami *et al.*, 1998; de Kloet & de Kloet, 2004; Henriques *et al.*, 2010). Within all of these genomes we identified three conserved amino acid motifs (FTLNN, GTPHLQGY and YCSK), a P-loop for dNTP binding (GPPGCGKS, amino acids 165 to 172) and three additional conserved motifs (WWDGY, amino acids 190 to 194; DDFYGWLP, amino acids 203 to 210; DRYP, amino acids 219 to 222) within the *rep* gene (Henriques *et al.*, 2010), and the nuclear localization domain (RRRYARPxYRR RHIRRYRLRRRRHFRRRRFxTNRIYTLRxKRQ) on the N-terminus of the CP (Heath *et al.*, 2006).

Within the full genome of all Italian strains, ORF3, ORF4, ORF5 and ORF7 displayed a high degree of homology when compared with the corresponding sequences reported by Bassami *et al.* (1998) and Henriques *et al.* (2010). ORF3 and ORF5 were detected in all isolates. ORF4 and ORF7, truncated or incomplete, were considered absent only in three strains (1IT08, 9IT11, 10IT11). ORF6 was absent in all strains analysed. For details, see Table 5.



**Figure 1.** Bone marrow (BM) histological sections: aPF and CF of PBFD. CF bone marrow (1a), bone marrow in aPF is partially replaced by fat tissue (atrophy) (1b). In the same sections, immunohistochemical staining (BFDV antigen): a lower number of BFDV-positive myeloid cells are present in CF (1c) compared with aPF (arrows) (1d). Numerous BFDV-positive cells are also positive for caspase-1 (arrows) (1e, 1f). A similar pattern of nuclear positivity is observed in caspase-1 and TUNEL stained myeloid cells (1g, 1h), with a higher number of TUNEL-positive cells being present in the aPF, particularly concentrated near the areas of adipose-metaplasia of the tissue (1h). Scale bar: 250  $\mu$ m.

**Table 3.** Immunohistochemical and TUNEL results on 10 African grey parrots (AGPs) necropsied in this study. Archived formalin-fixed, paraffin wax-embedded tissues from 3 AGPs deceased for accidental causes, negative for BFDV, were used as control.

Organ	Group	caspase-1	IL-1 $\beta$	TUNEL	BFDV
Thymus	aPF <sup>a</sup>	42.3%	11.25	38.2%	13%
	CF <sup>b</sup>	27.5%	5	17%	8%
	EP <sup>c</sup>	2%	1.5	0.5%	0%
Bursa	aPF	52.18%	11.5	42.8%	28%
	CF	37%	3	35.55%	15%
	EP	7%	0.5	4%	0%
Spleen	aPF	61%	10.75	51.2%	31%
	CF	44.5%	8.5	48.5%	27%
	EP	13.5%	1.5	11.5%	0%
Bone marrow	aPF	55.8%	12.25	46.3%	35.7%
	CF	4.5%	2.5	3%	2.5%
	EP	2%	0	1.5%	0%

<sup>a</sup>aPF (atypical peracute form: 8 parrots); <sup>b</sup>CF (chronic form: 2 parrots); <sup>c</sup>EP (euthanized parrots: 3 parrots).

The number of BFDV, caspase-1, and T.U.N.E.L.-positive cells is normalized to the number of cells per field and express as a percentage of these values.

The total score of IL-1 $\beta$  positive cells results from the mean value obtained from the scores deriving from each organs examined per all the bird of each category (aPF, CF, EP).

**Phylogenetic analysis.** The phylogenetic analysis of the 10 complete nucleotide sequences of BFDV strains isolated in Italy from CAGPs clustered in subtypes of the BFDV-J strain. Moreover, according to the subdivision into subgroups proposed by Varsani *et al.* (2011), seven samples were included into subgroups J1 (2-3-4-5-6-7IT09, 8IT10) and one into J<sub>2</sub> (9IT11). Furthermore, in agreement with these criteria (pairwise identity and phylogenetic distribution), we proposed two new subtypes: J<sub>6</sub> (10IT11) and J<sub>7</sub> (1IT08) (Figure 2). Notably, the pairwise sequence identity between subtypes ranged from 98 to 100%.

The full-length genomes of Italian viruses clustered into two close major clades belonging to the BFDV-J strain, together with other isolates from Germany (PEG07), Poland (PL573; PL688; PL785), Portugal (PT08; PT08-2&3, PT09, PT09-2) and the UK (PEU01), confirming that BFDV-J is the main strain infecting CAGPs in Europe, as described previously (Varsani *et al.*, 2011; Julian *et al.*, 2013). Two strains from Portugal and some strains from Poland (I and T subtypes) clustered with strains from CAGPs from Thailand and South Africa, belonging to other subtypes (Figure 2).

The phylogenetic analysis of the deduced amino acid coat protein (Figure 3) showed that 1IT08 together with the partial coat sequence 20IT11 clustered in the subtype M1. Thus, we decided to evaluate possible recombination events. This analysis showed a recombination event in the *cp* gene of 1IT08 (complete sequence) and in the *cp* gene of 20IT11. In detail, the 1IT08 displayed a recombination between M and F subtypes, while 20IT11 a recombination with M and I subtypes. The pairwise identity of the complete 1IT08 genome in comparison with the other J strain was 93.6%, slightly under the threshold of 94% that was considered by Varsani *et al.* (2011) as the BFDV strain demarcation. Considering the pairwise identity, the phylo-genetic distribution and the relevant genomic features of BFDV strains, we decided to classify the 1IT08 as a subtype J<sub>7</sub>.

In other samples, the phylogenetic analysis of CP (Figure 3) and REP (data not shown) proteins reflects the division into two major subgroups, namely J1 and J2, as suggested by the analysis of the full nucleotide sequence. We could not observe significant differences comparing sequences isolated from birds affected by aPF and sequences of viruses isolated in parrots displaying CF of the illness, either within our isolates or including sequences already present in the literature. Moreover, we did not observe any correlation between the phylogenetic distribution and regional origin of the samples, since strains from northern Italy (5IT09, 7IT09, 9IT11) grouped together with others from central Italy (2-3-4-6IT09, 8IT10). As expected, the sequences from the same breeding centre clustered most closely among them (>99.4% coat nucleotide pairwise identity).

## Discussion

To gather more information concerning the virus causing PBFD in Italy, we have characterized the strains originating from outbreaks of aPF beak and feather disease in young CAGPs that were fledged and weaned. The aPFs described in this study were characterized by sudden death,

or general good bodily condition without any classical feather or beak alterations but with severe leukopenia and anaemia. This was associated with a high incidence of apoptosis in the lymphoid organs and especially in the BM. In addition, apoptotic caspase-1-positive and TUNEL-positive cells also showed concurrent positivity to BFDV staining without the typical botryoid and basophilic inclusion bodies. This particular clinical and post-mortem presentation of the disease is very similar to those reported by Schoemaker *et al.* (2000) in young CAGPs.

We analysed both nucleotide and amino acid sequences of the *cp* genes from 20 birds, and in 10 of these (eight from young birds with aPF and two from adults with CF of the disease) necropsy was performed and the complete genome characterized. The main genomic features of these strains were compared with BFDV sequences available on the GenBank database. BFDV from CAGPs isolated in Italy clustered in subtypes of the J-strain: this finding is in agreement with the literature that ascribes the J strains as characteristic of European countries (Varsani *et al.*, 2011; Julian *et al.*, 2013). Moreover, according to the classification criteria proposed by Varsani *et al.* (2011) we suggested two new subtypes: J<sub>6</sub> and J<sub>7</sub>. Of interest, two birds showed that novel recombination events had taken place among BFDV strains belonging to the subtypes M, I and F, these strains being commonly present in Africa and Indonesia, which are the two major areas from which parrots imported into Italy originate (Bert *et al.*, 2005). We observed that the viral sequences of strains that produce the peracute illness clustered closely with the strains responsible for the chronic disease, thus suggesting the absence of a specific BFDV genetic variant accountable for a specific form of PBFD, as formerly hypothesized (Ypelaar *et al.*, 1999; Bassami *et al.*, 2001; Heath *et al.*, 2004; Raue *et al.*, 2004; Hughes & Piontkivska, 2008). On the basis of this observation, we have hypothesized that the clinical and pathological differences between aPF and CF of the disease could be strictly related to the age and host immune system status, rather than genomic differences between viral strains. The parrots with the aPF in this study were severely anaemic. Since much evidence supports the view that the main sites of virus replication are the haemocyctoblasts in the BM and the precursor lymphocytes in the lymphoid organs (Adair, 2000), this anaemia could result from the direct effect of the virus on the erythroblast precursor cells in the BM. Chicken anaemia virus, a member of the *Circoviridae* family, affects T-cell precursors during the development of the thymic cortex, leading to depletion of helper (CD4) and cytotoxic (CD8) T-cell populations in chicks (Randall *et al.*, 1984; Goryo *et al.*, 1987; Otaki *et al.*, 1988; Adair *et al.*, 1991; Bounous *et al.*, 1995; Adair, 1996, 2000; Adair & McNulty, 1997). Since we observed extensive lymphoid depletion, it would appear that BFDV, similarly to chicken anaemia virus, has an immunosuppressive effect in young parrots. Here we show that BFDV induces different apoptosis phenomena in juvenile birds compared with adults. Until now, the mechanism of BFDV-induced apoptosis was unknown. A possible mechanism could be the direct effect of a BFDV-encoded protein on virus infected and uninfected bystander cells, as in the apoptin encoded by chicken anaemia virus, a 14 kDa proline-rich protein that causes p53-independent but caspase-3 -dependent apoptosis in thymic lymphoblasts, intra-sinusoidal and extra-sinusoidal haemocyctoblasts, and reticular cells (Zhuang *et al.*, 1995a, b; Danen-van Oorschot *et al.*, 1999, 2000; Pietersen & Noteborn, 2000). Even if we cannot exclude non-specific phagocytosis, the frequent co-localization of viral antigens and apoptotic events we observed is suggestive of a direct association between BFDV infection and apoptosis. The fact that porcine circovirus 2 (a close relative of BFDV) activates caspase-8 and caspase-3 pathways, leading to apoptosis in the lymphoid organs (Liu *et al.*, 2005), lends support to this hypothesis. The severity of apoptosis in porcine circovirus 2 is also age dependent, with the most severe occurring in the youngest pigs. In contrast to the other caspases, caspase-1 is generally believed to be pro-inflammatory (Ghayur *et al.*, 1997; Thornberry & Lazebnik, 1998). Caspase-1 (also referred to as IL-1 $\beta$  converting enzyme) regulates lipopoly-saccharide-induced interferon- $\gamma$  production and in macrophages leads to cleavage of the precursor IL-1 $\beta$  into active IL-1 that subsequently initiates an intense host inflammatory response. In this study, the strong up-regulation of caspase-1 in young parrots clinically affected by aPF, corresponded with an up-regulation of IL-1  $\beta$ . This would account for the inflammation characterized by the infiltration of heterophils, plasma cells and macrophages that was observed. As described in porcine circovirus 2 infected pigs (Liu *et al.*, 2005), ORF3 of BFDV may play a major role in the age-related levels of induction of apoptosis and inflammation through activation of the caspase-1/IL-1 $\beta$  pathway, by the synthesis of a specific apoptin-like protein. As the tropism of BFDV for rapidly dividing cells is known (Todd *et al.*, 2001), we can speculate that young parrots, with their faster growth rate, are more sensitive to BFDV infection. This is in agreement with the higher percentage of BFDV-positive cells that we observed in aPF affected parrots when compared with CF-positive birds. Consequently, the clinical and pathological differences observed in the aPF and the CF of the disease could ultimately be due to the different

activation of caspase-1, IL-1 and apoptotic pathways due to different amounts of BFDV present.

In conclusion, the data presented support the hypothesis that, in the absence of a defined BFDV genetic variant accountable for a specific clinical form of PBF, differences between aPF and CF are strictly host related.

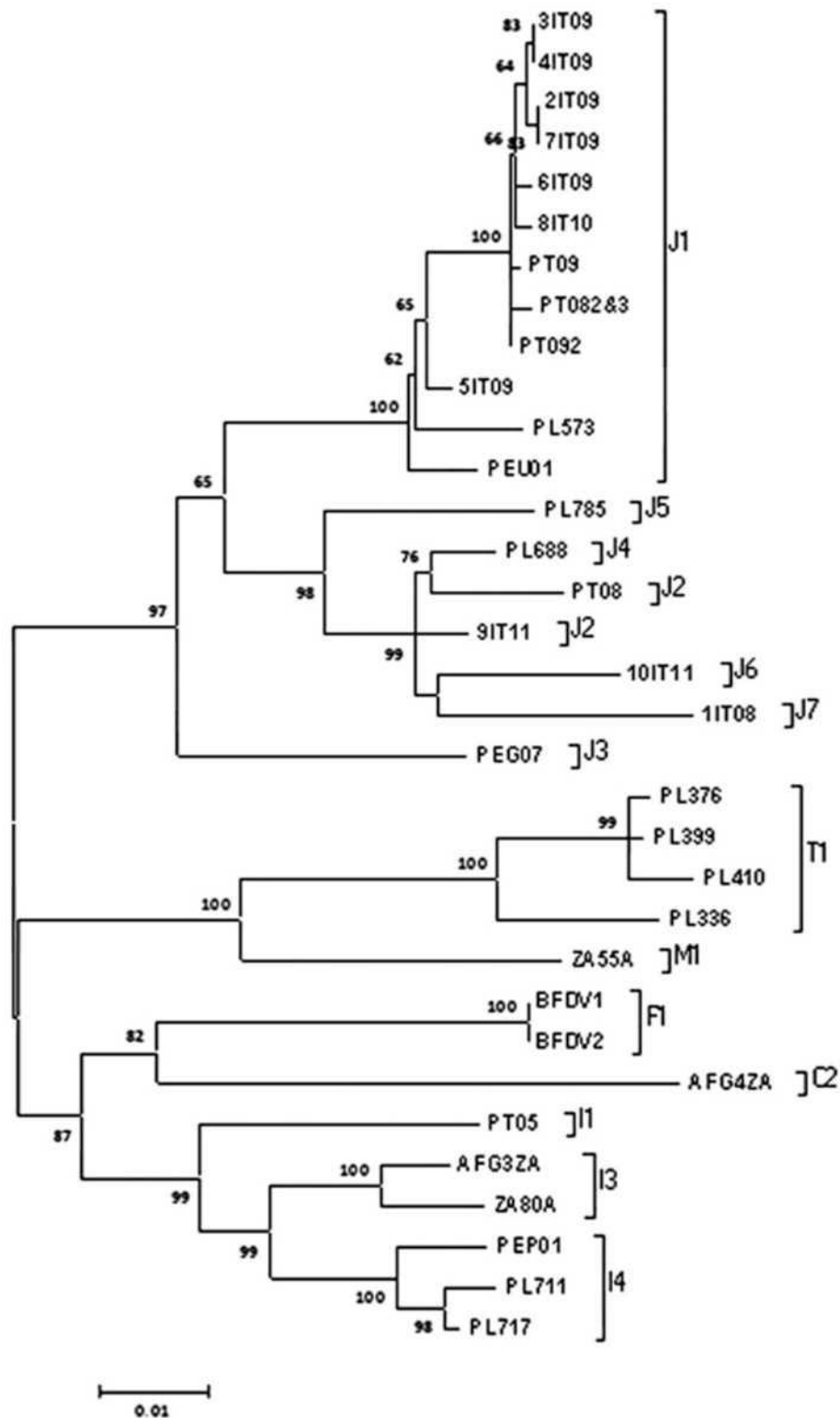
**Table 4.** Genomic features of the PBFV strains isolated in Italy (reference sequence: PEU01, GenBank accession AY521238).

Genomic features		3-4-5-7IT09; 8IT10	2IT09; 6IT09	1IT08	9IT11	10IT11
		location (nt)				
Intergenic region	76 nt	1986-2000 and 1-61	1987-2001 and 1-61	1984-1998 and 1-61	1985-1999 and 1-61	(75 nt) 1983-1997 and 1-60
Stem structure	CCGCCGCC	3-10	3-10	-	-	-
	GGCGGCGG	1985-1992	1986-1993	-	-	-
	CCGCCGCCT	-	-	3-11	3-11	3-11
	AGGCGGCGG	-	-	1982-1990	1983-1991	1981-1989
Octamer repeats and [internal hexamers]	G[GGGCAC]C	12-19 20-27	12-19 20-27	12-19 20-27	12-19 20-27	12-19 20-27
	G[GTGCCC]C	43-50 1974-1981	43-50 1975-1982	- 1972-1979	- 1973-1980	43-50 1971-1978
Only hexamer	GTGCCC	-	-	52-57	52-57	51-56
TATA boxes	TATA	86-89	86-89	-	-	-
		938-941	938-941	-	-	934-937
		1749-1752 <sup>a</sup>	-	1747-1750	1748-1751	1746-1749
	TATAAA	680-685	680-685	677-682	677-682	676-681
Polyadenylation signals	AATAAA	1197-1202	1198-1203	1195-1200	1196-1201	1194-1199
		-	-	-	-	1503-1508
		1237-1232	1238-1233	1235-1230	1236-1231	1234-1229
		1461-1456	1462-1457	-	-	-

<sup>a</sup> only 5IT09

**Table 5.** List and properties of the ORFs detected in the PBFDV strains isolated in Italy (reference sequence: PEU01, GenBank accession AY521238).

Strains			ORF1	ORF 2	ORF 3	ORF 4	ORF 5	ORF 6	ORF 7
<b>2IT09</b>	Location	nt	131-1000	1986-1237	559-62		543-1013		1301-1017
	Gene length	nt	870	750	498	Not detected	471	Not detected	285
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	165 (18.3)		156 (17)		94 (9.7)
<b>3IT09</b>	Location	nt	131-1000	1985-1236	559-62	1050-1388	543-1013		1300-1034
	Gene length	nt	870	750	498	339	471	Not detected	267
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	165 (18.3)	112 (11.9)	156 (17)		88 (9.0)
<b>4IT09</b>	Location	nt	131-1000	1985-1236	559-62	1050-1388	543-1013		1300-1034
	Gene length	nt	870	750	498	339	471	Not detected	267
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	165 (18.3)	112 (11.9)	156 (17)		88 (9.0)
<b>5IT09</b>	Location	nt	131-1000	1985-1236	559-62	1050-1388	543-1013		1300-1034
	Gene length	nt	870	750	498	339	471	Not detected	267
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	165 (18.3)	112 (11.9)	156 (17)		88 (9.0)
<b>6IT09</b>	Location	nt	131-1000	1986-1237	559-62		543-1013		1301-1017
	Gene length	nt	870	750	498	Not detected	471	Not detected	285
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	165 (18.3)		156 (17)		94 (9.7)
<b>7IT09</b>	Location	nt	131-1000	1985-1236	559-62	1050-1388	543-1013		1300-1034
	Gene length	nt	870	750	498	339	471	Not detected	267
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	165 (18.3)	112 (11.9)	156 (17)		88 (9.0)
<b>8IT10</b>	Location	nt	131-1000	1985-1236	559-62			Not detected	1300-1034
	Gene length	nt	870	750	498	339	Not detected	Not detected	267
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	165 (18.3)	112 (11.9)			88 (9.0)
<b>1IT08</b>	Location	nt	128-997	1983-1234	556-62		654-1016		1298- 1089
	Gene length	nt	870	750	495	Not detected	363	Not detected	210
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	164 (18.1)		120 (12.9)		69 (7.8)
<b>9IT11</b>	Location	nt	128-997	1984-1235	556-62		654-1016		
	Gene length	nt	870	750	495	Not detected	363	Not detected	Not detected
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	164 (18.1)		120 (12.9)		
<b>10IT11</b>	Location	nt	127-996	1982-1233	555-61		653-1015		1297- 1088
	Gene length	nt	870	750	495	Not detected	363	Not detected	210
	Protein length	aa (kDa)	289 (33.5)	249 (29.3)	164 (18.1)		120 (12.9)		69 (7.8)



**Figure 2.** Bayesian phylogenetic tree of the complete nucleotide sequences of BFDV obtained from the African grey parrots: 10 Italian strains and 14 BFDV NCBI reference sequences with strain and subtype demarcation. Posterior probability values are reported below branches. Bar: 0.01 substitutions per site. The phylogeny divided the BFDV into clusters and the sequences analysed in this study are included in group J and subgroups J1 to J7.





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